- ¹ Title: Tree height and leaf drought tolerance traits shape growth responses across droughts in a temperate
- 2 broadleaf forest
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22 Summary

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- As climate change is driving increased drought frequency and severity in many forested regions around the world, mechanistic understanding of the factors conferring drought resistance in trees is increasingly important. The dendrochronological record provides a window through which we can understand how tree size and species' traits shape tree growth responses during droughts.
- We analyzed tree-ring records for twelve species that comprise 97% of the woody productivity of the 25.6-ha ForestGEO plot in a broadleaf deciduous forest of northern Virginia (USA) to test hypotheses on how tree height, microenvironment characteristics, and species' traits shaped drought responses across the three strongest regional droughts over a 60-year period (1950 2009).
 - Individual-level drought resistance decreased with tree height, which was strongly correlated with
 exposure to higher evaporative demand and solar radiation. The potentially greater rooting volume
 of larger trees did not confer an advantage in sites with low topographic wetness index. Resistance
 was greater among species whose leaves experienced less shrinkage upon desiccation and lost turgor
 (wilted) at more negative water potentials.
- We conclude that tree height and leaf drought tolerance traits influence growth responses during
 drought, as recorded in the tree-ring record spanning historical droughts. Thus, these factors can be
 useful for predicting future drought responses under climate change.
- Key words: annual growth; crown exposure; drought; Forest Global Earth Observatory (ForestGEO); leaf drought tolerance traits; temperate broadleaf deciduous forest; tree height; tree-ring

41 Introduction

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Forests play a critical global role in climate regulation (Bonan, 2008), yet there remains enormous
   uncertainty as to how the forest-dominated terrestrial carbon sink will respond to climate change
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   (Friedlingstein et al., 2006). An important aspect of this uncertainty lies with physiological responses of
   trees to drought (Kennedy et al., 2019). In many forested regions around the world, the risk of severe
   drought is increasing (Trenberth et al., 2014; Dai et al., 2018), often despite increasing precipitation
   (Intergovernmental Panel on Climate Change, 2015; Cook et al., 2015). Droughts, intensified by climate
   change, have been affecting forests worldwide and are expected to continue as one of the most important
   drivers of forest change in the future (Allen et al., 2010, 2015). Understanding forest responses to drought
   requires elucidation of how tree size, microenvironment, and species' traits jointly influence individual-level
   drought resistance, defined here as a tree's ability to maintain growth during drought, and the extent to
   which their influence is consistent across droughts. Because the resistance and resilience of growth to
   drought is linked to trees' probability of surviving drought (DeSoto et al., 2020; Liu et al., 2019),
   understanding growth responses can also help elucidate which trees are most vulnerable to drought-induced
   mortality. However, it has proven difficult to resolve the many factors affecting tree growth during drought
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   with available forest census data, which only rarely captures extreme drought, and with tree-ring records,
   which capture multiple droughts but usually only sample a subset of a forest community, typically focusing
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   on a single species or the largest individuals.
   Many studies have shown that within and across species, large trees tend to be more affected by drought.
   Greater growth reductions for larger trees were first shown on a global scale by Bennett et al. (2015), and
   subsequent studies have reinforced this finding (e.g., Hacket-Pain et al., 2016; ?). It has yet to be resolved
   which of several potential underlying mechanisms most strongly shape these trends in drought response.
62
   First, tree height itself may be a primary driver. Taller trees face the biophysical challenge of lifting water
   greater distances against the effects of gravity and friction (McDowell et al., 2011; McDowell and Allen,
   2015; Ryan et al., 2006; Couvreur et al., 2018). Vertical gradients in stem and leaf traits-including smaller
   and thicker leaves (higher leaf mass per area, LMA), greater resistance to hydraulic dysfunction (i.e., more
   negative water potential at 50% loss of hydraulic conductivity, more negative P50), and lower hydraulic
   conductivity at greater heights (Couvreur et al., 2018; Koike et al., 2001; McDowell et al., 2011)-enable
   trees to become tall (Couvreur et al., 2018). Greater stem capacitance (i.e., water storage capacity) of
   larger trees may also confer resistance to transient droughts [Phillips et al. (2003)*; Scholz et al. (2011)].
   Indeed, tall trees require xylem of greater hydraulic efficiency, such that xylem conduit diameters are wider
   in the basal portions of taller trees, both within and across species (Olson et al., 2018; Liu et al., 2019),
   and throughout the conductive systems of angiosperms (Zach et al., 2010; Olson et al., 2014, 2018). Wider
   xylem conduits plausibly make large trees more vulnerable to embolism during drought (Olson et al.,
   2018), and traits conducive to efficient water transport may also lead to poor ability to recover from or
   re-route water around embolisms (Roskilly et al., 2019).
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   Larger trees may also have lower drought resistance because of microenvironmental and ecological factors.
77
   Their crowns tend to occupy more exposed canopy positions, which are associated with higher evaporative
   demand (Kunert et al., 2017). Subcanopy trees tend to fare better specifically due to the benefits of a
   buffered environment (Pretzsch et al., 2018). Counteracting the liabilities associated with tall height, large
   trees tend to have larger root systems (Enquist and Niklas, 2002), potentially mitigating some of the
   biophysical challenges they face by allowing greater access to water. Larger root systems—if they grant
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access to deeper water sources—would be particularly advantageous in drier microenvironments (e.g.,
    hilltops, as compared to valleys and streambeds) during drought. Finally, tree size-related responses to
    drought can be modified by species' traits and their distribution across size classes (Meakem et al., 2018;
    Liu et al., 2019). Understanding the mechanisms driving the greater relative growth reductions of larger
    trees during drought requires sorting out the interactive effects of height and associated exposure, root
    water access, and species' traits.
    Debates have also arisen regarding the traits influencing tree growth responses to drought. Studies within
    temperate broadleaf forests have observed ring-porous species showing higher drought tolerance than
90
    diffuse-porous species (Friedrichs et al., 2009; Elliott et al., 2015; Kannenberg et al., 2019), but this
    distinction would not hold in the global context (Wheeler et al., 2007; Olson et al., 2020) and does not
    resolve differences among the many species within each category. Commonly-measured traits including
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    wood density and leaf mass per area (LMA) have been linked to drought responses within some temperate
    deciduous forests (Abrams, 1990; Guerfel et al., 2009; Hoffmann et al., 2011; Martin-Benito and Pederson,
    2015) and across forests worldwide (Greenwood et al., 2017). However, in other cases these traits could not
    explain drought tolerance (e.g., in a tropical rainforest; Maréchaux et al., 2019), or the direction of
97
    response was not always consistent. For instance, higher wood density has been associated with greater
    drought resistance at a global scale (Greenwood et al., 2017), but correlated negatively with tree
    performance during drought in a broadleaf deciduous forest in the southeastern United States (Hoffmann
100
    et al., 2011). Thus, the perceived influence of these traits on drought resistance may actually reflect
101
    indirect correlations with other traits that more directly drive drought responses (Hoffmann et al., 2011).
102
    In contrast, hydraulic traits have direct physiological linkages to tree growth and mortality responses to
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    drought. For instance, water potentials at which percent the loss of conductivity surpasses a certain
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    threshold (e.g., P50 and P88, representing 50 and 88% loss of conductivity, respectively) and hydraulic
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    safety margin (i.e., difference between typical minimum water potentials and P50 or P88) correlate with
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    drought performance across global forests (Anderegg et al., 2016). However, these are time-consuming to
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    measure and therefore infeasible for predicting or modeling drought responses in highly diverse forests
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    (e.q., in the tropics). More easily-measurable leaf drought tolerance traits that have direct linkage to plant
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    hydraulic function can explain variation in plant distribution and function (Medeiros et al., 2019). These
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    include leaf area shrinkage upon desiccation (PLA_{dry}; Scoffoni et al., 2014) and the leaf water potential at
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    turgor loss point (\pi_{tlp}), i.e., the water potential at which leaf wilting occurs (Bartlett et al., 2016a; Zhu
    et al., 2018). Both traits correlate with hydraulic vulnerability and drought tolerance as part of unified
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    plant hydraulic systems (Scoffoni et al., 2014; Bartlett et al., 2016a; Zhu et al., 2018; Farrell et al., 2017).
114
    The abilities of both PLA_{dry} and \pi_{tlp} to explain tree drought resistance remains untested.
115
    Here, we examine how tree height, microenvironment characteristics, and species' traits collectively shape
    drought resistance, defined as the ratio of annual growth in a drought year to that which would be
117
    expected in the absence of drought based on previous years' growth. We test a series of hypotheses and
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    associated specific predictions (Table 1) based on the combination of tree-ring records from the three
    strongest droughts over a 60-year period (1950 - 2009), species trait measurements, and census and
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    microenvironmental data from a large forest dynamics plot in Virginia, USA. First, we focus on how tree
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    size, alone and in its interaction with microenvironmental gradients, influences drought resistance. We
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    examine the contemporary relationship between tree height and microenvironment, including growing
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    season meteorological conditions and crown exposure. We then test whether, consistent with most forests
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globally, larger-diameter, taller trees tend to have lower drought resistance in this forest, which is in a region (eastern North America) represented by only two studies in the global review of Bennett et al. (2015). We also test for an influence of potential access to available soil water, which should be greater 127 for larger trees in dry but not in perpetually wet microsites. Finally, we focus on the role of species' traits, 128 testing the hypothesis that species' traits-particularly leaf drought tolerance traits-predict drought resistance. We test predictions that drought resistance is higher in ring-porous than semi-ring and 130 diffuse-porous species and that it is correlated with wood density-either positively (Greenwood et al., 2017) 131 or negatively (Hoffmann et al., 2011) and positively correlated with LMA. We further test predictions that species with low PLA_{dry} have higher drought resistance, and that species whose leaves lose turgor lower 133 water potentials (more negative π_{tlp}) have higher resistance. 134

135 Materials and Methods

- 136 Study site and microclimate
- 137 Research was conducted at the 25.6-ha ForestGEO (Forest Global Earth Observatory) study plot at the
- Smithsonian Conservation Biology Institute (SCBI) in Virginia, USA (38°53'36.6"N, 78°08'43.4"W; Fig.
- 139 S1) (Bourg et al., 2013; Anderson-Teixeira et al., 2015a). SCBI is located in the central Appalachian
- 140 Mountains near the northern boundary of Shenandoah National Park. Elevations range from 273 to 338 m
- above sea level with a topographic relief of 65m (Bourg et al., 2013). Climate is humid temperate, with
- mean annual temperature of 12.7°C and precipitation of 1005 mm yr⁻¹ during our study period (1960-2009;
- source: CRU TS v.4.01; Harris et al., 2014). Dominant tree taxa within this secondary forest include
- Liriodendron tulipifera, oaks (Quercus spp.), and hickories (Carya spp.; Table 2).
- $_{145}$ Identifying drought years
- We identified the three largest droughts within the time period 1950-2009, defining drought (Slette et al...
- ¹⁴⁷ 2019) as events with anomalously dry peak growing season climatic conditions. Specifically, we used the
- metric of Palmer Drought Severity Index (PDSI) during May-August (MJJA; Table S1), which were
- identified by Helcoski et al. (2019) as the months of the current year to which annual tree growth was most
- 150 sensitive at this site. PDSI divisional data for Northern Virginia were obtained from NOAA
- 151 (https://www7.ncdc.noaa.gov/CDO/CDODivisionalSelect.jsp) in December 2017. Based on this, we
- 152 identified the three strongest droughts during the study period (Figs. 1, S1; Table S1).
- The droughts differed in intensity and antecedent moisture conditions (Fig. S1, Table S1). The 1966
- drought was preceded by two years of moderate drought during the growing season and severe to extreme
- drought starting the previous fall. In August 1966, PDSI reached its lowest monthly value (-4.82) of the
- three droughts. The 1977 drought was the least intense throughout the growing season, and it was
- preceded by 2.5 years of near-normal conditions, making it the mildest of the three droughts. The 1999
- drought was preceded by wetter than average conditions until the previous June, but PDSI plummeted
- below -3.0 in October 1998 and remained below this threshold through August 1999.
- 160 Data collection and preparation
- Within or just outside the ForestGEO plot, we collected data on a suite of variables including tree heights,
- microenvironment characteristics, and species traits (Table 3). The SCBI ForestGEO plot was censused in
- $_{163}$ 2008, 2013, and 2018 following standard ForestGEO protocols, whereby all free-standing woody stems \geq

(Condit, 1998). From these census data, we used measurements of DBH from 2008 to calculate historical DBH and data for all stems > 10cm to analyze functional trait composition relative to tree height (all 166 analyses described below). Census data are available through the ForestGEO data portal 167 (www.forestgeo.si.edu). We analyzed tree-ring data (xylem growth increment) from 571 trees representing the twelve dominant species (Table 2; Fig. S2). Selected species were those with the greatest contributions to woody 170 aboveground net primary productivity $(ANPP_{stem})$ and together comprised 97% of study plot $ANPP_{stem}$ 171 between 2008 and 2013 (Helcoski et al., 2019). Cores (one per tree) were collected within the ForestGEO plot at breast height (1.3m) in 2010-2011 or 2016-2017. In 2010-2011, cores were collected from randomly 173 selected live trees of each species that had at least 30 individuals ≥ 10 cm DBH (Bourg et al., 2013). 174 Annual tree mortality censuses were initiated in 2014 (Gonzalez-Akre et al., 2016), and in 2016-2017, cores were collected from all trees found to have died since the previous year's census. We note that drought was 176 probably not a cause of mortality for these trees, as monthly May-Aug PDSI did not drop below -1.75 in 177 these years or the three years prior (2013-2017), and that trees cored dead displayed similar climate sensitivity to trees cored live (Helcoski et al., 2019). Cores were sanded, measured, and crossdated using 179 standard procedures, as detailed in (Helcoski et al., 2019). The resulting chronologies (Fig. 1a) were 180 published in Zenodo (DOI: 10.5281/zenodo.2649302) in association with Helcoski et al. (2019). For each cored tree, we combined tree-ring records and allometric equations of bark thickness to reconstruct DBH for the years 1950-2009. Prior DBH was estimated using the following equation:

1cm diameter at breast height (DBH) were mapped, tagged, measured at DBH, and identified to species

$$DBH_Y = DBH_{2008} - 2 * \left[r_{bark,2008} - r_{bark,Y} + \sum_{year=Y}^{2008} r_{ring,Y} \right]$$

Here, Y denotes the year of interest, r_{ring} denotes ring width derived from cores, and r_{bark} denotes bark thickness. Bark thickness was estimated from species-specific allometries based on the bark thickness data

from the site (Anderson-Teixeira et al., 2015b). Specifically, we used linear regression on log-transformed 186 data to relate r_{bark} to diameter inside bark from 2008 data (Table S2), which were then used to determine 187 r_{bark} in the DBH reconstruction. Tree heights (H) were measured by several researchers for a variety of purposes between 2012 and 2019 (n=1,518 trees). Methods included direct measurements using a collapsible measurement rod on small trees (NEON, 2018) or a tape measure on recently fallen trees (this study); geometric calculations using 191 clinometer and tape measure (Stovall et al., 2018a) or digital rangefinders (Anderson-Teixeira et al., 2015b; NEON, 2018); and ground-based LiDAR (Stovall et al., 2018b). Rangefinders used either the tangent 193 method (Impulse 200LR, TruPulse 360R) or the sine method (Nikon ForestryPro) for calculating heights. 194 Both methods are associated with some error (Larjavaara and Muller-Landau, 2013), but in this instance there was no clear advantage of one or the other. Measurements from the National Ecological Observatory Network (NEON) were collected near the ForestGEO plot following standard NEON protocol, whereby 197 vegetation of short stature was measured with a collapsible measurement rod, and taller trees with a rangefinder (NEON, 2018). Species-specific height allometries were developed (Table S3) using log-log 199 regression $(ln[H] \sim ln[DBH])$. For species with insufficient height data to create reliable species-specific 200 allometries (n=2, JUNI and FRAM), heights were calculated from an equation developed by combining the

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year, Y, based on reconstructed DBH_Y. The distribution of H across drought years is shown in Fig. S3.
    To characterize how environmental conditions vary with height, data were obtained from the NEON tower
    located <1km from the study area via the neonUtilities package (Lunch et al., 2020). We used wind speed.
205
    relative humidity, and air temperature data, all measured over a vertical profile spanning heights from 7.2
206
    m to above the top of the tree canopy (31.0 or 51.8m, depending on censor), for the years 2016-2018
    (NEON, 2018). After filtering for missing and outlier values, we determined the daily minima and maxima.
208
    which we then aggregated at the monthly scale.
209
    Crown position—a categorical variable classifying trees based on exposure to sunlight—was recorded for all
210
    cored trees that remained standing during the growing season of 2018 following the protocol of Jennings
211
    et al. (1999). Trees were classified as follows: dominant trees were defined as those with crowns above the
212
    general level of the canopy, co-dominant trees as those with crowns within the canopy; intermediate
213
    trees as those with crowns below the canopy level, but illuminated from above; and suppressed as those
    below the canopy and receiving minimal direct illumination from above.
215
    Topographic wetness index (TWI), used here as a metric of long-term mean moisture availability, was
216
    calculated using the dynatopmodel package in R (Fig. S2) (Metcalfe et al., 2018). Originally developed by
    Beven and Kirkby (1979), TWI was part of a hydrological run-off model and has since been used for a
218
    number of purposes in hydrology and ecology (Sørensen et al., 2006). TWI calculation depends on an input
219
    of a digital elevation model (DEM; ~3.7 m resolution from the elevatr package (Hollister, 2018)), and from
    this yields a quantitative assessment defined by how "wet" an area is, based on areas where run-off is more
221
    likely. From our observations in the plot, TWI performed better at categorizing wet areas than the
222
    Euclidean distance from the stream.
223
    Species' trait data were collected in August 2018 (Tables 2-3; Fig. S4). We sampled small, sun-exposed
224
    branches up to eight meters above the ground from three individuals of each species in and around the
225
    ForestGEO plot. Sampled branches were re-cut under water at least two nodes above the original cut and
226
    re-hydrated overnight in covered buckets under opaque plastic bags before measurements were taken.
227
    Rehydrated leaves taken towards the apical end of the branch (n=3 per individual: small, medium, and
228
    large) were scanned, weighed, dried at 60° C for ≥ 48 hours, and then re-scanned and weighed. Leaf area
229
    was calculated from scanned images using the LeafArea R package (Katabuchi, 2019). LMA was
230
    calculated as the ratio of leaf dry mass to fresh area. PLA_{dry} was calculated as the percent loss of area
231
    between fresh and dry leaves. Wood density was calculated for ~1cm diameter stem samples (bark and pith
232
    removed) as the ratio of dry weight to fresh volume, which was estimated using Archimedes' displacement.
233
    We used the rapid determination method of Bartlett et al. (2012) to estimate osmotic potential at turgor
234
    loss point (\pi_{tlp}). Briefly, two 4 mm diameter leaf discs were cut from each leaf, tightly wrapped in foil,
235
    submerged in liquid nitrogen, perforated 10-15 times with a dissection needle, and then measured using a
    vapour pressure osmometer (VAPRO 5520, Wescor, Logan, UT, USA). Osmotic potential (\pi_{osm}) given by
237
   the osmometer was used to estimate (\pi_{tlp}) using the equation \pi_{tlp} = 0.832\pi_{osm}^{-0.631} (Bartlett et al., 2012).
238
    Statistical Analysis
239
    For each drought year, we calculated a metric drought resistance (Rt) as the ratio of basal area increment
240
    (BAI; i.e., change in cross-sectional area) during the drought year to the mean BAI over the five years
241
    preceding the drought (Lloret et al., 2011). Thus, Rt values <1 and >1 indicate growth reductions and
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height measurements across all species. We then used these allometries to estimate H for each drought

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increases, respectively. Because the Rt metric could be biased by directional pre-drought growth trends, we
243
    also tried an intervention time series analysis (ARIMA, (Hyndman et al., 2020)) that predicted mean
    drought-year growth based on trends over the preceding ten years and used this value in place of the
245
    five-year mean in calculations of resistance (Rt_{ARIMA}= observed BAI/ predicted BAI). The two metrics
246
    were strongly correlated (Fig. S5). Visual review of the individual tree-ring sequences with the largest
    discrepancies between these metrics revealed that Rt was less prone to unreasonable estimates than
248
    Rt_{ARIMA}, so we selected Rt as our focal metric, presenting parallel results for Rt_{ARIMA} in the
249
    Supplementary Info. In this study we focus exclusively on drought resistance metrics (Rt or Rt_{ARIMA}),
    and not on the resilience metrics described in Lloret et al. (2011), because (1) we would expect resilience to
251
    be controlled by a different set of mechanisms, and (2) the findings of (DeSoto et al., 2020) suggest that Rt
252
    is a more important drought response metric for angiosperms in that low resistance to moderate droughts
253
    was a better predictor of mortality during subsequent severe droughts than the resilience metrics.
254
    Analyses focused on testing the predictions presented in Table 1 with Rt as the response variable, and then
255
    repeated using Rt_{ARIMA} as the response variable. Models were run for all drought years combined and for
256
    each drought year individually. The general statistical model for hypothesis testing was a mixed effects
257
    model, implemented in the lme4 package in R (Bates et al., 2019). In the multi-year model, we included a
258
    random effect of tree nested within species and a fixed effect of drought year to represent the combined
259
    effects of differences in drought characteristics. Individual year models included a random effect of species.
    All models included fixed effects of independent variables of interest (Tables 1,3) as specified below. All
261
    variables across all best models had variance inflation factors <1.2 (1 + /- 0.019). We used AICc to assess
262
    model selection, and conditional/marginal R-squared to assess model fit as implemented in the
263
    AICcmodavg package in R (Mazerolle and portions of code contributed by Dan Linden., 2019). AICc refers
264
    to a corrected version of AICc, and is best suited for small data sizes (see Brewer et al., 2016).
265
    To avoid over-fitting models with five species traits (Table 3) across only 12 species, we did not include all
266
    traits as fixed effects in a single linear mixed model, but rather conducted individual tests of each species
267
    trait to determine the relative importance and appropriateness for inclusion in the main model. These tests
268
    followed the model structure specified above, then added ln[H] and ln[TWI] to create a base model
269
    against which we tested traits. Trait variables were considered appropriate for inclusion in the main model
270
    if they had a consistent direction of response across all droughts and if their addition to the base model
271
    improved fit (at \triangle AICc \ge 1.0) in at least one drought year (Table S4). We note that we did not use the
    \DeltaAICc \geq 1.0 criterion as a test of significance, but rather of whether the variable had enough influence to
273
    be considered as a candidate variable in full models.
274
    We then determined the top full models for predicting Rt (or Rt_{ARIMA}). To do so, we compared models
275
    with all possible combinations of candidate variables, including ln[H]*ln[TWI] and species traits as
    specified above. We identified the full set of models within \triangle AICc=2 of the best model (that with lowest
277
    AICc). When a variable appeared in all of these models and the sign of the coefficient was consistent across
278
    models, we viewed this as support for the acceptance/rejection of the associated prediction (Table 1). If
    the variable appeared in some but not all of these models, and its sign was consistent across models, we
280
    considered this partial support/rejection. In presentation of the results below, we note instances where the
281
    Rt_{ARIMA} model disagreed with the Rt model, but otherwise do not discuss the Rt_{ARIMA} model.
282
    All analysis beyond basic data collection was performed using R version 3.6.2 (R Core Team, 2019). Other
283
    R-packages used in analyses are listed in the Supplementary Information (Appendix S1). All data, code,
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- and results are available through the SCBI-ForestGEO organization on GitHub
- 286 (https://github.com/SCBI-ForestGEO: SCBI-ForestGEO-Data and
- ²⁸⁷ McGregor_climate-sensitivity-variation repositories), with static versions corresponding to data and
- ²⁸⁸ analyses presented here archived in Zenodo (DOIs: 10.5281/zenodo.3604993 and [TBD], respectively.

289 Results

- 290 Tree height and microenvironment
- 291 In the years for which we have vertical profiles in climate data (2016-2018), taller trees—or those in
- dominant crown positions—were generally exposed to higher evaporative demand during the peak growing
- season months (May-August; Fig. 2). Specifically, maximum daily wind speeds were significantly higher
- above the top of the canopy (40-50m) than within and below (10-30m) (Fig. 2a). Relative humidity was
- also somewhat lower during June-August, ranging from $\sim 50-80\%$ above the canopy and $\sim 60-90\%$ in the
- understory (Fig. 2b). Air temperature did not vary consistently across the vertical profile (Fig. 2c).
- ²⁹⁷ Crown position varied as expected with height (dominant > co-dominant > intermediate > suppressed),
- but with substantial variation (Fig. 2d). There were significant differences in height across all crown
- position classes (Fig. 2d). A comparison test between height and crown position data from the most recent
- ForestGEO census (2018) revealed a correlation of 0.73.
- 301 Community-level drought responses
- At the community level, cored trees showed substantial growth reductions in all three droughts, with a
- $_{303}$ mean Rt of 0.86 in 1966 and 1999, and 0.84 in 1977 (Fig. 2b). Across the entire study period (1950-2009),
- the focal drought years were the three years with the largest fraction of trees exhibiting Rt < 0.7.
- Specifically, in each drought, roughly 30% of the cored trees had growth reductions of $\geq 30\%$ ($Rt \leq 0.7$):
- ³⁰⁶ 29% in 1966, 32% in 1977, and 27% in 1999. However, some individuals exhibited increased growth, i.e.,
- Rt > 1.0: 26% of trees in 1966, 22% in 1977, and 26% in 1999.
- In the context of the multivariate model, Rt did not vary across drought years. That is, drought year as a
- variable did not appear in any of the top models -i.e., models that were statistically indistinguishable
- $(\Delta AICc < 2)$ from the best model.
- 311 Tree height, microenvironment, and drought resistance
- Taller trees (based on H in the drought year) showed stronger growth reductions during drought (Table 1;
- Figs. 4, S6). Specifically, ln[H] appeared, with a negative coefficient, in the best model ((Δ AICc=0) and
- all top models when evaluating the three drought years together (Tables S6-S7). The same held true for
- ³¹⁵ 1966 individually. For the 1977 drought, ln[H] did not appear in the best model, but was included, with a
- negative coefficient, among the top models-i.e., models that were statistically indistinguishable
- $(\Delta AICc < 2)$ from the best model (Tables 1, S6-S7). For the 1999 drought, ln[H] had no significant effect.
- Rt had a significantly negative response to ln[TWI] across all drought years combined (Figs. 4, S6, Table
- S6-S7). The effect was also significant for 1977 and 1999 individually (Fig. 4, Table S6). When Rt_{ARIMA}
- was used as the response variable, the effect was significant in 1977, and included in some of the top
- models in 1966 and 1999 (Table S7). This negates the idea that trees in moist microsites would be less
- affected by drought. Nevertheless, we tested for a ln[H] * ln[TWI] interaction, a negative sign of which

in drier microenvironments with a deeper water table. This hypothesis was rejected, as the ln[H] * ln[TWI] interaction was never significant, and had a positive sign in any top models in which it 325 appeared (Tables 1, S6-S7). This term did appear with a positive coefficient in the best Rt_{ARIMA} model 326 for all years combined (Table S7), indicating that the negative effect of height on Rt was significantly stronger in wetter microhabitats. 328 Species' traits and drought resistance 329 Species, as a factor in ANOVA, had significant influence (p<0.05) on all traits (wood density, LMA, 330 PLA_{dry} , and π_{tlp}), with more significant pairwise differences for wood density and PLA_{dry} than for LMAand π_{tlp} (Table 2, Fig. S4). Drought resistance also varied across species, overall and in each drought year 332 (Fig. 3). Significant differences in Rt across species were most pronounced in 1966 with a total of seven 333 distinct groupings, while 1977 had four and 1999 had two. Averaged across all droughts, Rt was lowest in Liriodendron tulipifera (mean Rt = 0.66) and highest in Fagus grandifolia (mean Rt = 0.99). 335 Wood density, LMA, and xylem porosity were all poor predictors of Rt (Tables 1,S4-S5). Wood density 336 and LMA were never significantly associated with Rt in the single-variable tests and were therefore 337 excluded from the full models. Xylem porosity was also excluded from the full models, as it had no significant influence for all droughts combined and had contrasting effects in the individual droughts: 339 whereas ring-porous species had higher Rt than diffuse- and semi-ring- porous species in the 1966 and 1999 340 droughts, they had lower Rt in 1977 (Table S4). It is noteworthy that the two diffuse-porous species in our study, Liriodendron tulipifera and Fagus grandifolia, were at opposite ends of the Rt spectrum (Fig. 3), 342 further refuting the idea that xylem porosity is a useful predictor of Rt in the context of this study. 343 In contrast, PLA_{dry} , and π_{tlp} were both negatively correlated to drought resistance (Figs. 4, S6; Tables 344 1,S4-S7). Both had consistent signs across all droughts, and their inclusion at least marginally improved 345 the model ($\triangle AICc > 1.0$) for at least one of the three droughts (Table S4), qualifying them as candidate 346 variables for the full model. PLA_{dry} had a significant influence, with negative coefficient, in full models for 347 the three droughts combined and for the 1966 drought individually (Fig. 4; Tables S6-S7). For 1977 and 348 1999, it was included with a negative coefficient in some of the top models (Tables S6-S7). π_{tlp} was included 349 with a negative coefficient in the best model for both all droughts combined and for the 1977 drought 350 individually (Fig. 4; Table S6). It was also included in some of the top models for 1999 (Tables S6-S7). 351

could indicate that smaller trees (presumably with smaller rooting volume) are more susceptible to drought

2 Discussion

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Tree height, microenvironment, and leaf drought tolerance traits shaped tree growth responses across three 353 droughts at our study site (Table 1, Fig. 4). The greater susceptibility of larger trees to drought, similar to 354 forests worldwide (Bennett et al., 2015), was driven primarily by their height (Stovall et al., 2019). Taller 355 height was likely a liability in itself, and was also associated with greater exposure to conditions that would 356 promote water loss and heat damage during drought (Fig. 2). There was no evidence that greater 357 availability of, or access to, soil water availability increased drought resistance; in contrast, trees in wetter 358 topographic positions had lower Rt (Zuleta et al., 2017; Stovall et al., 2019), and the larger potential rooting volume of large trees provided no advantage in the drier microenvironments. The negative effect of 360 height on Rt held after accounting for species' traits, which is consistent with recent work finding height 361 had a stronger influence on mortality risk than forest type during drought (Stovall et al., 2020). Drought

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resistance was not consistently linked to species' LMA, wood density, or xylem type (ring- vs. diffuse
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    porous), but was negatively correlated with leaf drought tolerance traits (PLA_{dry}, \pi_{tlp}). This is the first
    study to our knowledge linking PLA_{dry} and \pi_{tlp} to growth reduction during drought. The directions of
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    these responses were consistent across droughts (Table S6), supporting the premise that they were driven
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    by fundamental physiological mechanisms. However, the strengths of each predictor varied across droughts
    (Fig. 4; Tables S6-S7), indicating that drought characteristics interact with tree size, microenvironment,
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    and traits to shape which individuals are most affected. These findings advance our knowledge of the
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    factors that make trees vulnerable to growth declines during drought and, by extension, likely make them
    more vulnerable to mortality (Sapes et al., 2019).
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    The droughts considered here were of a magnitude that has occurred with an average frequency of
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    approximately once every 10-15 years (Fig. 1a, Helcoski et al., 2019) and had substantial but not
    devastating impacts on tree growth (Figs. 1b). These droughts were classified as severe (PDSI < -3.0;
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    1977) or extreme (PDSI < -4.0; 1966, 1999) at our site and have been linked to tree mortality in the
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    eastern United States (Druckenbrod et al., 2019). However, extreme, multiannual droughts such as the
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    so-called "megadroughts" of this type that have triggered massive tree die-off in other regions (e.g., Allen
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    et al., 2010; Stovall et al., 2019) have not occurred in the Eastern United States within the past several
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    decades (Clark et al., 2016). Of the droughts considered here, the 1966 drought, which was preceded by
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    two years of dry conditions (Fig. S1), severely stressed a larger portion of trees (Fig. 1b). The tendency
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    for large trees to have lowest resistance was most pronounced in this drought, consistent with other
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    findings that this physiological response increases with drought severity (Bennett et al., 2015; Stovall et al.,
    2019). Across all three droughts, the majority of trees experienced reduced growth, but a substantial
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    portion had increased growth (Fig. 1b), consistent with prior observations that smaller trees can exhibit
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    increased growth rates during drought (Bennett et al., 2015). It is likely because of the moderate impact of
    these droughts, along with other factors influencing tree growth (e.g., stand dynamics), that our best
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    models characterize only a modest amount of variation in Rt: 11-12% for all droughts combined, and
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    18-25% for each individual drought (Fig. S6; Table S6).
    Consistent with studies in other forests worldwide (Bennett et al., 2015), taller trees in this forest exhibited
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    lower drought resistance. Mechanistically, this is consistent with, and reinforces, previous findings that
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    biophysical constraints make it impossible for trees to efficiently transport water to great heights and
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    simultaneously maintain strong resistance and resilience to drought-induced embolism (Olson et al., 2018;
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    Couvreur et al., 2018; Roskilly et al., 2019). Taller trees also face dramatically different microenvironments
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    (Fig. 2). They are exposed to higher wind speeds and lower humidity (Fig. 2a-b), resulting in higher
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    evaporative demand. Unlike other temperate forests where modestly cooler understory conditions have
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    been documented (Zellweger et al., 2019), particularly under drier conditions (Davis et al., 2019), we
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    observed no significant variation in air temperatures across the vertical profile (Fig. 2c). More critically for
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    tree physiology, leaf temperatures can become significantly elevated over air temperature under conditions
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    of high solar radiation and low stomatal conductance (Campbell and Norman, 1998; Rey-Sánchez et al.,
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    2016). Under drought, when air temperatures tend to be warmer, direct solar radiation tends to be higher
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    (because of less cloud cover), and less water is available for evaporative cooling of the leaves, trees with
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    sun-exposed crowns may not be able to simultaneously maintain leaf temperatures below damaging
    extremes and avoid drought-induced embolism. Indeed, previous studies have shown lower drought
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    resistance in more exposed trees (Liu and Muller, 1993; Suarez et al., 2004; Scharnweber et al., 2019).
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    Unfortunately, collinearity between height and crown exposure in this study (Fig. 2d) makes it impossible
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and mature forest stands, along with short and tall isolated trees, would be valuable for more clearly disentangling the roles of tree height and crown exposure. 408 Belowground, taller trees would tend to have larger root systems (Enquist and Niklas, 2002), but this does 409 not necessarily imply that they have greater access to or reliance on deep soil-water resources that may be 410 critical during drought. While tree size can correlate with the depth of water extraction (?), the linkage is 411 not consistent. Shorter trees can vary broadly in the depth of water uptake (?), and larger trees may 412 allocate more to abundant shallow roots that are beneficial for taking up water from rainstorms (Meinzer et al., 1999). Moreover, reliance on deep soil-water resources can actually prove a liability during severe 414 and prolonged drought, as these can experience more intense water scarcity relative to non-drought 415 conditions (?). In any case, the potentially greater access to water did not override the disadvantage conferred by height—and, in fact, greater moisture access in non-drought years (here, higher TWI) appears 417 to make trees more sensitive to drought (Zuleta et al., 2017; Stovall et al., 2019). This may be because 418 moister habitats would tend to support species and individuals with more mesophytic traits (Bartlett 419 et al., 2016b; Mencuccini, 2003; Medeiros et al., 2019), potentially growing to greater heights (e.g., Detto 420 et al. (2013)), and these are then more vulnerable when drought hits. The observed height-sensitivity of 421 Rt, together with the lack of conferred advantage to large stature in drier topographic positions, agrees 422 with the concept that physiological limitations to transpiration under drought shift from soil water 423 availability to the plant-atmosphere interface as forests age (Bretfeld et al., 2018), such that tall, dominant 424 trees are the most sensitive in mature forests. Again, additional research comparing drought responses across forests with different tree heights and water availability would be valuable for disentangling the 426 relative importance of above- and belowground mechanisms across trees of different size. 427 The development of tree-ring chronologies for the twelve most dominant tree species at our site (Helcoski et al., 2019; Bourg et al., 2013) gave us the sample size to compare historical drought responses across 429 species (Fig. 3) and associated traits at a single site (see also Elliott et al., 2015). Our study reinforced 430 current understanding (see Introduction) that wood density and LMA are not reliably linked to drought resistance (Table 1). Contrary to previous studies in temperate deciduous forests, we did not find an 432 association between xylem porosity and drought resistance, as the two diffuse-porous species, Liriodendron 433 tulipifera and Faqus grandifolia, were at opposite ends of the Rt spectrum (Fig. 3). While the low Rt of L. 434 tulipifera is consistent with other studies (Elliott et al., 2015), the high Rt of F. grandifolia contrasts with 435 studies identifying diffuse porous species in general (Elliott et al., 2015; Kannenberg et al., 2019), and the 436 genus Faqus in particular (Friedrichs et al., 2009), as drought sensitive. There are two potential 437 explanations for this discrepancy. First, other traits can and do override the influence of xylem porosity on 438 drought resistance. Ring-porous species are restricted mainly to temperate deciduous forests, while highly 439 drought-tolerant diffuse-porous species exist in other biomes (Wheeler et al., 2007). Fagus grandifolia had 440 intermediate π_{tlp} and low PLA_{dry} (Fig. S4), which would have contributed to its drought resistance (Fig. 441 4; see discussion below). A second explanation of why F. grandifolia trees at this particular site had higher 442 Rt is that the sampled individuals, reflective of the population within the plot, are generally shorter and in 443 less-dominant canopy positions compared to most other species (Fig. S4). The species, which is highly 444 shade-tolerant, also has deep crowns (Anderson-Teixeira et al., 2015b), implying that a lower proportion of leaves would be affected by harsher microclimatic conditions at the top of the canopy under drought (Fig. 446 2). Thus, the high Rt of the sampled F. grandifolia population can be explained by a combination of fairly 447 drought-resistant leaf traits, shorter stature, and a buffered microenvironment.

to confidently partition causality. Additional research comparing drought responses of early successional

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Concerted measurement of tree-rings and leaf drought tolerance traits of emerging importance (Scoffoni
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    et al., 2014; Bartlett et al., 2016a; Medeiros et al., 2019) allowed novel insights into the role of drought
    tolerance traits in shaping drought response. The finding that PLA_{dry} and \pi_{tlp} can be useful for predicting
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    drought responses of tree growth (Fig. 4; Table 1) is both novel and consistent with previous studies
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    linking these traits to habitat and drought tolerance. Previous studies have demonstrated that \pi_{tlp} and
    PLA_{dry} are physiologically meaningful traits linked to species distribution along moisture gradients
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    (Maréchaux et al., 2015; Fletcher et al., 2018; Medeiros et al., 2019; Simeone et al., 2019; Rosas et al., 2019;
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    Zhu et al., 2018), and our findings indicate that these traits also influence drought responses. Furthermore,
    the observed linkage of \pi_{tlp} to Rt in this forest aligns with observations in the Amazon that \pi_{tlp} is higher
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    in drought-intolerant than drought-tolerant plant functional type. Further, it adds support to the idea that
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    this trait is useful for categorizing and representing species' drought responses in models (Powell et al.,
    2017). Because both PLA_{dry} and \pi_{tlp} can be measured relatively easily (Bartlett et al., 2012; Scoffoni
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    et al., 2014), they hold promise for predicting drought growth responses across diverse forests. The
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    importance of predicting drought responses from species traits increases with tree species diversity; whereas
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    it is feasible to study drought responses for all dominant species in most boreal and temperate forests (e.g.,
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    this study), this becomes difficult to impossible for species that do not form annual rings, and for diverse
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    tropical forests. Although progress is being made for the tropics (Schöngart et al., 2017), a full linkage of
    drought tolerance traits to drought responses would be invaluable for forecasting how little-known species
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    and whole forests will respond to future droughts (Christoffersen et al., 2016; Powell et al., 2017).
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    As climate change drives increasing drought in many of the world's forests (Trenberth et al., 2014;
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    Intergovernmental Panel on Climate Change, 2015), the fate of forests and their climate feedbacks will be
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    shaped by the biophysical and physiological drivers observed here. Our results, consistent with other
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    observations around the world, imply that the tallest, most exposed trees will be most affected (Bennett
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    et al., 2015; Stovall et al., 2019). We show that, at least within the mature forest studied here, the
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    vulnerability conferred by tall height and associated crown exposure outweigh any advantage of a larger
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    root system, even in drier microenvironments. This would suggest that the drought responses of trees in
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    mature forests are more strongly differentiated along the size spectrum by their above- than below-ground
    environment. The same may not be true of systems where short trees exist outside of a buffered understory
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    environment-i.e., open grown trees or short-statured, early-successional forests. The latter appear to be
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    limited more strongly by root water access during drought (Bretfeld et al., 2018), and would also be
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    dominated by species with different traits. The earlier-successional species at our site (Liriodendron
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    tulipifera, Quercus spp., Fraxinus americana) display a mix of traits conferring drought resistance (Table
480
    2), while the late-successional Fagus grandifolia displayed high drought resistance, in part because it exists
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    primarily within a buffered microenvironment. This has potentially far-reaching implications for
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    future forests. If we accept that climate change will cause more intense droughts, it is
    possible we will see a shift from mature forests with buffering, tall trees to forests with a
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    shorter overall height (e.g. see McDowell et al. (2020)). At this point, species that had high
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    resistance due to buffering (such as F. grandifolia) may in turn become more susceptible.
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    This shift, in addition to the susceptibility of large trees to fire and pests/pathogens, could be
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    highly detrimental to temperate forests. That being said, Further research on how leaf drought
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    tolerance traits and drought vulnerability change over the course of succession would be valuable for
    addressing how drought resistance changes as forests age (e.g. Rodríguez-Catón et al., 2015). In the
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    meantime, the results of this study advance our knowledge of the factors conferring drought resistance in a
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mature forest, opening the door for more accurate forecasting of forest responses to future drought.

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504 Author Contribution

- KAT, IM, and AJT designed the research. Tree-ring chronologies were developed by RH under guidance of
- 506 AJT and NP. Trait data were collected by IM, JZ under guidance of NK and LS. Other plot data were
- collected by IM, AS, EGA, and NB under guidance of EGA and WM. Data analyses were performed by IM
- under guidance of KAT and VH. KAT and IM interpreted the results. IM and KAT wrote the first draft of
- manuscript, and all authors contributed to revisions.

510 Supplementary Information

- $_{511}$ Table S1: Monthly Palmer Drought Severity Index (PDSI), and its rank among all years between 1950 and
- ⁵¹² 2009 (driest=1), for focal droughts.
- Table S2: Species-specific bark thickness regression equations.
- Table S3: Species-specific height regression equations.
- Table S4. Individual tests of species traits as drivers of drought resistance, where Rt is used as the
- 516 response variable.
- Table S5. Individual tests of species traits as drivers of drought resistance, where Rt_{ARIMA} is used as the
- ⁵¹⁸ response variable.
- Table S6. Summary of top full models for each drought instance, where Rt is used as the response variable.
- Table S7. Summary of top models for each drought instance, where Rt_{ARIMA} is used as the response
- variable.
- 522 Figure S1. Time series of Palmer Drought Severity Index (PDSI) for the 2.5 years prior to each focal
- $_{523}$ drought
- Figure S2: Map of ForestGEO plot showing topographic wetness index and location of cored trees
- Figure S3: Distribution of reconstructed tree heights across drought years.

- Figure S4. Distribution of independent variables by species.
- Figure S5. Comparison of Rt and Rt_{ARIMA} results, with residuals, for each drought scenario
- Figure S6. Visualization of best model, with data, for all droughts combined.

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